

PRIMERS, METHODS AND KITS FOR DETECTING KILLER-CELL IMMUNOGLOBULIN-LIKE RECEPTOR ALLELES

FIELD OF INVENTION

Embodiments of the present invention relate to primer pairs,
5 primer sets and methods of using the primer pairs and primer sets to
identify KIR alleles. Certain embodiments also encompass kits for use
with the present primers or methods. More particularly this invention
relates to primer pairs, primer sets and methods of identifying KIR alleles
by amplification of approximately 1000 bp amplicons from an intra-exon
10 portion of the KIR extracellular domain.

BACKGROUND OF THE INVENTION

Several types of highly diversified molecules, such as the
ABO blood group system, the family of MHC (Major Histocompatibility
Complex, called, in humans, HLA-human leukocyte antigen), the family of
15 receptors for the T lymphocyte antigen (TCR) and the family of receptors
for the B lymphocyte antigen (BCR) partially characterize immune
functions in humans and animals. Each of these different families of
molecules, when expressed by an adult individual, constitute an
individually specific repertoire that is involved in immune system self or
20 non-self recognition.

Immunologists have more recently identified other specific
immune system repertoires, such as the repertoire defined by natural
killer-cell immunoglobulin-like receptors (KIRs). The KIR family of natural
killer (NK) cell receptors, a family with currently approximately 14 genes
25 and 2 pseudogenes, is highly polymorphic and mimics on natural killer
cells the clonotypic expression of TCRs and BCRs on T-cells and B-cells.

The diverse members of the KIR family participate in mediating cell-cell recognition by NK cells.

Generally, NK cells exhibit cytotoxic activity following recognition of non-self during cell-cell interactions. Inhibitory KIR family members, whose natural ligands are members of the MHC Class I complex, prevent NK cytotoxicity upon ligand binding. Thus, when a KIR finds its natural ligand on a cell, it recognizes the cell as self. This mechanism suggests that inhibitory KIR receptors play a role in preventing autoimmune reactions. Inhibitory KIRs may also participate in the antigenic incompatibility during allograft or xenograft transplantation.

In in vivo allograft hemopoietic transplantation, scientists have demonstrated the involvement of KIRs in a graft versus leukemia, a positive side effect of allograft transplantation. In patients undergoing hemopoietic transplants, a balance must be achieved between the incidence of graft versus host disease, i.e. where the transplanted cells begin to attack the healthy cells of the recipient, and graft versus leukemia, i.e. where the transplanted cells only attack pathogenic cells. KIRs present on the graft but lacking a matched MHC Class I ligand in the recipient have been implicated in a reduced risk of leukemia relapse in patients receiving bone marrow transplants. Presumably, KIRs contribute to an increase in graft versus leukemia. Although scientists recognize the beneficial effects of graft versus leukemia, currently grafted tissue or cells are not screened for cells that contribute to graft versus leukemia. As KIRs appear to be involved in this process, it would be highly beneficial to be able to screen transplanted tissue or cells in order to determine KIRs that may increase graft versus leukemia.

Effectively typing individual alleles of KIRs in a graft or in a patient is complicated by the degree of complexity in KIR expression. The currently available means used in the medical context do not make it possible to easily and inexpensively document all the presently known KIR

repertoires during transplantation. Only the compatibility of the HLA-A, HLA-B, and HLA-DR molecules of the donor and of the recipient are currently checked prior to a grant or transplant. Although systems have been developed in an attempt to establish KIR typing, none of the
5 presently available systems provide a way to type all presently known KIR alleles. Because KIR alleles are closely related, their close homology often makes it impossible to discriminate between alleles without sophisticated procedures such as nucleotide sequencing.

Thus, there continues to be a need for identifying known KIR
10 receptors.

SUMMARY OF THE INVENTION

In one embodiment a primer pair for identifying a killer-cell immunoglobulin-like receptor allele is described. The primer set consists of primers capable of amplifying all presently known KIR alleles.

15 According to this primer set amplicons that are less than or 1000 bases in length are amplified from an intra-exon portion of a nucleic acid that encodes for an extracellular portion of a KIR receptor.

In alternative embodiments, amplicon size may vary and amplicons may be less than or 500 or 250 bases in length or greater than
20 or 2000 bases in length.

Based on these primer sets, methods of detecting KIR alleles using the primer sets are described. Kits for carrying out these methods are also provided in some embodiments. These kits can include instructions for carrying out the methods, one or more reagents useful in
25 carrying out these methods, and one or more primer sets capable of amplifying all presently known KIR alleles.

Objects and advantages of the present invention will become more readily apparent from the following detailed description.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is an image of an electrophoretic gel showing the results of amplifying a sample containing KIR alleles with a primer set described herein with different thermal cyclers.

5 Figure 2 is an enlarged image of an electrophoretic gel showing the results of amplifying a sample containing KIR alleles with a primer set described herein.

 Figure 3 is an image of an electrophoretic gel showing the results of amplifying a sample containing KIR alleles with a primer set
10 described herein with different DNA template quantities.

 Figure 4 is an image of an electrophoretic gel showing the results of amplifying a sample containing KIR alleles with a primer set described herein with different DNA polymerase amounts.

 Figure 5 is an image of an electrophoretic gel showing the
15 results of amplifying a sample containing KIR alleles with a primer set described herein with a denaturing temperature in the thermal cycling reaction of 92°C.

 Figure 6 is an image of an electrophoretic gel showing the results of amplifying a sample containing KIR alleles with a primer set
20 described herein with an annealing temperature in the thermal cycling reaction of 61°C.

 Figure 7 is an image of an electrophoretic gel showing the results of amplifying a sample containing KIR alleles with a primer set described herein with an annealing temperature in the thermal cycling
25 reaction of 65°C.

 Figure 8 shows selected IHW panel sample results using a primer set and method described herein.

DETAILED DESCRIPTION

Described herein are primer sets, methods and kits for detecting one or more killer-cell immunoglobulin-like receptor (KIR) alleles. KIRs, members of the immunoglobulin (Ig) superfamily, mediate the function of natural killer cells in innate immune responses and play an important role in regulating Natural Killer (NK) cell activity. KIR receptors are found on the surface of human NK cells and some T-cell subsets. These receptors recognize Class I MHC molecules expressed on target cells and some of these receptors directly interact with polymorphic HLA-A, -B or -C determinants. For example, inhibitory KIRs use HLA Class I as ligands (Bw4 and Cw epitopes). This recognition helps determine whether the target should be lysed – if the KIR fails to recognize the appropriate ligand, the NK cell becomes cytotoxic.

Some pathogens and tumor infected cells evade the immune system by down-regulating HLA Class I molecules. When HLA Class I is downregulated, cytotoxic T cells no longer have the opportunity to recognize and react to peptides bound to HLA. The immune system can counteract the loss of immuncity caused by down-regulated MHC Class I through the KIR / NK mechanism. This phenomenon, which results in cytotoxicity when NK cells bind to non-self cells, has been described as the “missing-self hypothesis.” KIRs involved in the missing-self hypothesis are inhibitory KIRs. Additionally, KIRs can be activating. However, the ligand is currently unknown for activating KIRs.

KIR diversity in individuals is achieved through many factors, including: allelic variability, gene content; gene copy number and gene expression. Individual KIR genes may be exhibited more than once on one haplotype. Expression of KIRs in individuals varies significantly and individuals may exhibit between 8 and 16 genes/pseudogenes.

KIR receptors and alleles are described in the following references, which are hereby incorporated by references. Carrington M, Norman P. The KIR Gene Cluster: Bethesda MD: National (USA) Library of Medicine, NCBI, 2003 <http://www.ncbi.nlm.nih.gov>; Hsu K, Liu X, Selvakumar A, Mickelson E, O'Reilly R, Dupont B. The Journal of Immunology, 2002, 169:5118-5129; Yawata M, Yawata N, Abi-Rached L, Parham P. Critical Reviews in Immunology, 2002, 22:463-482; Gomez-Lozano and Vilches, Tissue Antigen 2002: 59:184-193; Killer cell immunoglobulin-like receptor (KIR) Nomenclature report, 2002 Steven G.E. Marsh, et al. Human Immunology 64: 648-654; Genotyping of human killer-cell immunoglobulin-like receptor genes by polymerase chain reaction with sequence specific primers: Diverse, Rapidly Evolving Receptors of Innate and Adaptive Immunity; Vilches, C, Parham, P, Annual Review in Immunology, 2002 20:217-251; A structural perspective on MHC class I recognition by killer cell immunoglobulin-like receptors; Boyington, J, Sun, P, Molecular Immunology, 2001 38: 1007-1021; The killer cell immunoglobulin-like receptor (KIR) genomic region: gene-order, haplotypes and allelic polymorphism; Hsu, K, Chida, S, Geraghty, D, Dupont, B, Immunological Reviews, 2002 190:40-52; and Structure and function of major histocompatibility complex (MHC) class I specific receptors expressed on human natural killer (NK) cells; Borrego, F, Kabat, J, et al. Molecular Immunology 2001 38:637-660. Sequences of the presently known KIR alleles are reported at the Immuno Polymorphism Database (IPD) website www.ebi.ac.uk/ipd/kir/ and the National Center for Biotechnology Information (NCBI) website – dbMHC, www.ncbi.nlm.nih.gov/mhc/.

One embodiment of a primer set provides a primer set that identifies all of the presently known KIR alleles. Such a primer set can include a plurality of primer pairs such that a majority of the primer pairs are capable of producing an amplicon that is less than or 1000 bases in

length from a nucleic acid that encodes a KIR receptor. Amplification of smaller PCR products is more efficient and tolerant of assay variables such as degraded DNA and reduced polymerase activity. Accordingly, the present primer sets, methods and kits exhibit a robustness allowing them to be used with minimal experimentation. In some embodiments, the majority of the primer pairs are capable of producing an amplicon that is less than or 1000 bases in length from an intra-exon portion of a nucleic acid that encodes for a portion of a KIR receptor. In these and other embodiments, one or more of the primer pairs of the majority of the primer pairs are capable of producing an amplicon that is less than or 1000 bases in length from a nucleic acid that encodes for an extracellular portion of a KIR receptor.

In another embodiment, a primer set includes a first primer and second primer that together are capable of producing an amplicon that is less than or 1000 bases in length from an intra-exon portion of a nucleic acid that encodes for an extracellular portion of a KIR receptor. Accordingly, in this embodiment a KIR primer pair targets intra-exon polymorphism such that the gene-specific amplicon sizes do not exceed 1000 bp. As above, amplification of smaller PCR products is more efficient and tolerant of assay variables such as degraded DNA and reduced polymerase activity so that the primer sets, the described methods, and kits, are more robust than previous compositions and methods for determining individual KIR alleles. The primer sets may also contain additional primer pairs that are specific for a desired KIR allele. For example, one or more of such primer pairs can be capable of producing an amplicon that is less than or 1000 bases in length from an intra-exon portion of a nucleic acid that encodes for an extracellular portion of one or more additional KIR receptors. In some embodiments, the primer set contains primer pairs that are capable of identifying all presently known KIR receptors. In these primer sets a majority of the

primer pairs in the primer set are capable of producing an amplicon that is less than or 1000 bases in length from an intra-exon portion of a nucleic acid that encodes for an extracellular portion of the KIR receptors.

5 In any of the above primer sets a majority of the primer pairs can produce an amplicon that is less than or 250 or 500 bases in length. In certain embodiments a majority of the primer pairs are capable of producing an amplicon from about 100, 150, 200 or 250 to about 250, 500, 750 or 1000 bases in length. However, the present primer sets do not require that all the primer pairs produce amplicons from KIR alleles
10 that are less than or 1000 bases in length. As such, one or more primer pairs can be capable of producing an amplicon that is greater than 1000 bases in length. Further, primer pairs are capable of producing an amplicon from an inter-exon of a nucleic acid that encodes for a portion of a KIR receptor.

15 In some embodiments, none of the primer pairs are capable of producing an amplicon greater than or 2000, 3000, 4000 or 5000 bases in length. In further embodiments, in place of a majority about 25%, 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 75%, 80%, 85%, 90% or 95% of the primer pairs in any of the primer sets described
20 herein can have the recited characteristics.

Suitable intra-exon or extracellular KIR domains that can be amplified by the present primer pairs include those encoded by any one of KIR exons 1-8, and in particular the polymorphic regions of these exons. Different primer pairs can produce amplicons from different KIR exons as
25 desired. As will be apparent to those skilled in the art, the exact sequence of the primers or amplicons produced are not critical to the present primer sets, methods or kits as long as the KIR allele(s) being tested for can be specifically identified. As such, the present primers will generally target KIR allele sequences that are unique to the specific allele
30 and distinguish one KIR allele from the others. It will also be apparent

that with some KIR alleles such distinguishing sequences can be found in many different portions of the KIR gene. Accordingly, the primer pairs typically target polymorphism in the extracellular domains of the KIR receptors based on the fact that more nonsynonymous mutation events
5 occur in these regions compared to the intra cellular and transmembrane regions of each gene. Therefore the primer sets, method and kits can provide high resolution or allele level genotyping methods that primarily target polymorphism in the extra cellular domains of the KIR genes.

The individual primers in the primer sets can be of any
10 length, for example ranging from 5 nucleotides to several hundred. Preferably, the primer oligonucleotides will have a length of greater than 10 nucleotides, and more preferably, a length of from 12-50 nucleotides, such as 12-25 or 15-20 nucleotides. The primer oligonucleotides can also be chosen to have a desired melting temperature, such as 40 to 80°C, 50
15 to 70°C, 55 to 65°C, or 60°C. The length of the primer is sufficient to permit the primer oligonucleotide to hybridize to the target molecule. The sequence of the primer oligonucleotide is selected such that it is complementary to a predetermined sequence of the target molecule. The 3' terminus of the primers in the primer sets are capable of being
20 extended by a nucleic acid polymerase under appropriate conditions. The present primers can be used in any method where nucleic acid primers find utility. For example, the primers are readily applicable to RT PCR of KIR mRNA for expression analysis because they may target exon regions. The present primer pairs can also be used individually to identify a single
25 KIR allele, as desired. The present primers can also be extended to, as yet, unknown KIR alleles.

One example of an assay where the present primer pairs find use include detection assays or methods for identifying a KIR allele in a sample having, or suspected of having, a KIR allele. In such an assay,
30 generally, the sample will be contacted with the primer set under

conditions such that the primer pair will amplify the KIR allele for which the primer pair is specific, if that allele is present in the sample. The presence or absence of the amplicon can then be determined or detected by standard techniques, such as separation techniques including
5 electrophoresis, chromatography (including HPLC and denaturing-HPLC), or the like. Exemplary techniques for performing these assays are described in the examples section. As will be recognized by the skilled artisan, the production of a specific amplicon will indicate the presence of a specific KIR allele in a sample. Accordingly, the presence or absence of
10 an amplicon can be correlated with the presence or absence of the specific KIR allele in the sample. The sample to be detected can be obtained from any suitable source or by any suitable technique.

Typically, the nucleic acid amplification or extension of the KIR alleles involves mixing a target nucleic acid with a "master mix"
15 containing the reaction components for performing the amplification reaction. This reaction mixture is then subjected to temperature conditions that allow for the amplification of the target KIR. The reaction components in the master mix can include a buffer which regulates the pH of the reaction mixture; one or more of the four deoxynucleotides
20 (dATP, dCTP, dGTP, dTTP - preferably present in equal concentrations), which provide the energy and nucleosides necessary for the synthesis of DNA; primers or primer pairs that bind to the DNA template in order to facilitate the initiation of DNA synthesis; and a DNA polymerase that adds the deoxynucleotides to the complementary DNA strand being
25 synthesized. The polymerase used in the present methods and kits is not particularly limited, and any suitable polymerase can be used. Examples of suitable polymerase include thermostable polymerase enzymes, such as the Taq polymerase. Preferred polymerases exhibit low error rates during strand synthesis.

A typical thermal cycling reaction has a temperature profile that involves an initial ramp up to a predetermined, target denaturation temperature which is high enough to separate the double-stranded target DNA into single strands. Generally, the target denaturation temperature of the thermal cycling reaction is approximately 91-97°C and the reaction is held at this temperature for a time period ranging between 20 seconds to two minutes. Then, the temperature of the reaction mixture is lowered to a target annealing temperature which allows the primers to anneal or hybridize to the single strands of DNA. Annealing temperatures can vary greatly depending upon the primers and target DNA used. Individual KIR alleles may exhibit individual or equivalent annealing temperatures. Generally, annealing temperatures range from 37°C to 55°C depending upon the application. Next, the temperature of the reaction mixture is raised to a target extension temperature to promote the synthesis of extension products. The extension temperature is generally held for approximately two minutes and occurs at a temperature range from 50°C to 72°C. This completes one cycle of the thermal cycling reaction. The next cycle then starts by raising the temperature of the reaction mixture to the denaturation temperature. Typically, the cycle is repeated 25 to 35 times to provide the desired quantity of DNA. As will be understood by the skilled artisan, the above description of the thermal cycling reaction is provided for illustration only, and accordingly, the temperatures, times and cycle number can vary depending upon the nature of the thermal cycling reaction and application.

The present assays and methods can be performed using a single primer pair specific for a single KIR allele or can use a set of primers that have specificity for more than one KIR allele. In some embodiments, different primer sets are contained within different amplification vessels, such as different wells of a multi-well plate, so that only a single primer set specific for a single KIR allele is present in an individual amplification

vessel. Such a configuration simplifies use and interpretation of the assay results. However, the present assays can use multiplex configurations where two, three or more primer pairs that are specific for different KIR alleles can be used in the same reaction vessel and one or more reaction vessels can be utilized. Amplification reactions using different primer pairs can be run in parallel, simultaneously or subsequent to one another, as desired. In some embodiments, all of the primer pairs required to identify all presently known KIR alleles can be contained within the same reaction vessel. In these multiplex assays, typically the primer pairs will be designed so that the resulting amplicons that are specific for a single KIR allele can be distinguished from one another. For example, the amplicons for the different KIR alleles can all have different lengths, or the primer pairs or amplicons can have distinct labels or be distinctly labeled.

In some embodiments, the present KIR primer sets, methods and kits can use a standard sequence specific primer technique. In a non-limiting example, twenty pre-aliquoted primer pairs, plus an optional internal control as well as an optional negative contamination control, can be used to identify all presently known KIR alleles. The methods and kits can also include a nucleic acid amplification buffer, with or without a polymerase, which in some instances will be aliquoted in per test volumes. The present assays can also use positive and negative controls to help verify results.

In some embodiments, the present primer sets, assays and kits can identify the presence of all presently known alleles of KIR genes: 2DL1, 2DL2, 2DL3, 2DL4, 2DL5, 2DP1, 2DS1, 2DS2, 2DS3, 2DS4, 2DS5, 3DL1, 3DL2, 3DL3, 3DP1, and 3DS1, as well as the more recently described variants such as the 2DS4 homolog KIR1D. Primer sets including primer pairs specific for one, two, three, four, five, six, seven, eight, nine, ten, eleven, twelve, thirteen or more KIR alleles can be designed for producing an amplicon that is less than or 1000 bases in

length from an intra-exon portion of a nucleic acid that encodes for an extracellular portion of a KIR receptor. Other primer pairs that produce amplicons in excess of 1000 bases in length can be used in conjunction, such as in parallel, with primer pairs producing the smaller amplicon. All primer mixes can target polymorphism in the extracellular domains of KIRs. All primer mixes can also contain a distinct internal control primer set to ensure proper assay performance. The assay can be completed in less than 3 hours when starting with genomic DNA and can follow the four procedure steps typical of the sequence specific priming (SSP) technology: (1) prepare a master mix with sample DNA, (2) dispense the master mix into the SSP mixes, (3) thermal cycle, and (4) analyze the amplicons by gel electrophoresis. The SSP assay can provide reliable and unambiguous detection of all presently known KIR alleles in a simple and rapid format.

The present primer sets, methods and kits can be used to define or identify KIR haplotypes, genotypes and polymorphic variation in an individual or in different populations. Some embodiments can also be used to identify KIR compatible and incompatible stem cell transplant donor-recipient pairs as well as study if KIR mismatching between donor and recipient correlates with KIR epitope mismatch predicted by HLA. Some embodiments can be used to determine the effect of KIR receptors on post-transplant complications. KIR has also been implicated in association with a wide variety of diseases including HIV, rheumatoid vasculitis, psoriatic arthritis, as well as playing a role in bone marrow transplantation. Regarding the association of KIR with bone marrow research and transplantation, NK Cells have been implicated in promotion of engraftment and mediation of graft versus leukemia (GVL). More recently, it has been suggested that the graft versus leukemia effects are a result of donor-recipient HLA epitope mismatching for KIR. Consistent with "missing self hypothesis" – if transplant recipient lacks HLA Class I

determinant recognized by the donor KIR, the NK cells to become cytotoxic towards tumor cells, reducing the risk of relapse. Therefore a mismatch of KIR ligands between host and donor may be preferable to promote GVL. Accordingly, the present primers, methods and kits can be used for research and clinical applications for any KIR associated disease, disorder, condition or phenomenon.

Any or all of the present primers can be labeled with a detectable moiety, if desired, to facilitate detection. When present, the detectable moiety is not particularly limited. Suitable examples of detectable labels include fluorescent molecules, beads, polymeric beads, fluorescent polymeric beads and molecular weight markers. Polymeric beads can be made of any suitable polymer including latex or polystyrene.

Certain embodiments also provide arrays of individual primers, primer pairs and primer sets that are contained within distinct, defined locations on a support. Each defined, distinct area of the array will typically have a plurality of the same primers. In some embodiments, the primers will be physically attached to the support in the defined location. The primers can also be contained within a well of the support. As used herein the term well is used solely for convenience and is not intended to be limiting. For example, a well can include any structure that serves to hold the nucleic acid primers in the defined, distinct area on the solid support. Non-limiting example of wells include depressions, grooves, walled surroundings and the like. In some of the arrays, the primers at different location can have the same probing regions or consist of the same molecule. This embodiment is useful when testing whether nucleic acids from variety of sources contain the same target sequences. The arrays can also have primers with one or different primer regions at different location within the array. This embodiment can be useful where nucleic acids from a single source are assayed for a variety of target sequences. Combinations of these array configurations are also provided

where some of the primers in the defined locations contain the same primer regions whereas other locations contain primers with primer regions that are specific for different targets.

Any suitable support can be used for the present arrays,
5 such as glass or plastic, either of which can be treated or untreated to help bind, or prevent adhesion of, individual primers, primer pairs or primer sets. In some embodiments, the support will be a multi-well plate so that the primers need not be bound to the support and can be free in solution. Such arrays can be used for automated or high volume assays
10 for target nucleic acid sequences.

Although the present primers generally utilize the five standard nucleotides (A, C, G, T and U) in their nucleotide sequences, the identity of the nucleotides or nucleic acids are not so limited. Non-standard nucleotides and nucleotide analogs, such as peptide nucleic
15 acids and locked nucleic acids can be used as desired. Several nucleotide analogs are known in the art (e.g., see, in Rawls, C & E News Jun. 2, 1997 page 35; in Brown, Molecular Biology LabFax, BIOS Scientific Publishers Limited; Information Press Ltd, Oxford, UK, 1991). In addition, the bases in a primer sequence may be joined by a linkage other than a
20 phosphodiester bond, so long as the bond does not interfere with hybridization with KIR alleles. Nucleotide analogs can include any of the known base analogs of DNA and RNA such as, but not limited to, 4-acetylcytosine, 8-hydroxy-N6-methyladenosine, aziridinylcytosine, pseudoisocytosine, 5-(carboxyhydroxymethyl) uracil, 5-fluorouracil, 5-bromouracil, 5-carboxymethylaminomethyl-2-thiouracil, 5-
25 carboxymethylaminomethyluracil, dihydrouracil, hypoxanthine, inosine, N6-isopentenyladenine, 1-methyladenine, 1-methylpseudouracil, 1-methylguanine, 1-methylinosine, 2,2-dimethylguanine, 2-methyladenine, 2-methylguanine, 3-methylcytosine, 5-methylcytosine, N6-methyladenine,
30 7-methylguanine, 5-methylaminomethyluracil, 5-methoxy-aminomethyl-2-

thiou- racil, beta-D-mannosylqueosine, 5'-methoxycarbonylmethyluracil, 5-methoxyuracil, 2-methylthio-N6-isopentenyladenine, uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid, oxybutoxosine, pseudouracil, queosine, 2-thiocytosine, 5-methyl-2-thiouracil, 2-thiouracil, 4-thiouracil, 5-methyluracil, N-uracil-5-oxyacetic acid methylester, uracil-5-oxyacetic acid, pseudouracil, queosine, 2-thiocytosine, orotic acid, 2,6-diaminopurine and the AEGIS™ bases isoC and isoG. As such, the individual primers, primer pairs and primer sets can contain DNA, RNA, analogs thereof or mixtures (chimeras) of these components.

Universal nucleotides can also be used in the present primers. As used herein, universal nucleotide, base, nucleoside or the like, refers to a molecule that can bind to two or more, i.e., 3, 4, or all 5, naturally occurring bases in a relatively indiscriminate or non-preferential manner. In some embodiments, the universal base can bind to all of the naturally occurring bases in this manner, such as 2'-deoxyinosine (inosine). For example, the universal base can bind all of the naturally occurring bases with equal affinity, such as 3-nitropyrrole 2'-deoxynucleoside (3-nitropyrrole) and those disclosed in U.S. Patent Nos. 5,438,131 and 5,681,947. Generally, when the base is "universal" for only a subset of the natural bases, that subset will generally either be purines (adenine or guanine) or pyrimidines (cytosine, thymine or uracil). Examples of nucleotides that can be considered universal for purines are known as the "K" base (N6-methoxy-2,6-diaminopurine), as discussed in Bergstrom et al., Nucleic Acids Res. 25:1935 (1997) and pyrimidines are known as the "P" base (6H,8H-3,4-dihydropyrimido[4,5-c][1,2]oxazin-7-one), as discussed in Bergstrom et al., supra, and U.S. Patent No. 6,313,286. Other suitable universal nucleotides include 5-nitroindole (5-nitroindole 2'-deoxynucleoside), 4-nitroindole (4-nitroindole 2'-deoxynucleoside), 6-nitroindole (6-nitroindole 2'-deoxynucleoside) or 2'-deoxynebularine. A partial order of duplex stability has been found as

follows: 5-nitroindole > 4-nitroindole > 6-nitroindole > 3-nitropyrrole. When used, such universal bases can be placed in polymorphic positions, for example those that are not required to specifically identify an allele. Combinations of these universal bases can also be used as desired.

5 Certain embodiments also provide kits for carrying out the methods described herein. In one embodiment, the kit is made up of one or more of the described primer pairs or primer sets with instructions for carrying out any of the methods described herein. The instructions can be provided in any intelligible form through a tangible medium, such as
10 printed on paper, computer readable media, or the like. A plurality of each primer pair or primer set can be provided in a separate container for easy aliquoting. The present kits can also include one or more reagents, buffers, hybridization media, salts, nucleic acids, controls, nucleotides, labels, molecular weight markers, enzymes, solid supports, dyes,
15 chromatography reagents and equipment and/or disposable lab equipment, such as multi-well plates (including 96 and 384 well plates), in order to readily facilitate implementation of the present methods. Such additional components can be packaged together or separately as desired. Solid supports can include beads and the like whereas molecular weight
20 markers can include conjugatable markers, for example biotin and streptavidin or the like. Enzymes that can be included in the present kits include DNA polymerases and the like. Examples of preferred kit components can be found in the description above and in the following examples.

25 In one embodiment of a kit, the kit contains sequence specific primers to identify the common forms of the following KIR genes: 2DL1, 2DL2, 2DL3, 3DL1, 3DL2, 2DS1, 2DS2, 2DS3, 2DS4, 2DS5, 3DS1, 2DL4, 2DL5, 3DL3, 3DP1, 2DP1. The primer mixes of the kit are present in trays that function under universal reaction conditions,
30 although reaction conditions can differ. Each kit contains 12 tests and 3

or 4 tests per tray are possible. Exemplary, but non-limiting, primer sets are described in Tables 2 and 3. This invention is further illustrated by the following non-limiting examples. These examples demonstrate the present primer sets, methods and kits provide reliable and unambiguous
5 detection of all presently known KIR genes in a simple, fast and cost-effective format.

EXAMPLES

For KIR genotyping of a single individual, the following protocol was used in the present examples, unless otherwise indicated.

10 **Step 1:** Make Mastermix solution (dNTP's, PCR-buffer, MgCl₂, Ficoll, Loading dye)

Step 2: Vortex thoroughly.

Step 3: Dispense 8 μ l of the Mastermix solution in each well. Add .12 μ l Taq Polymerase (5U/ μ l); 3 μ l water; 1.25 μ l DNA (at 75-
15 125 ng/ μ l) to the Mastermix solution. Total final reaction volume in each well is 13 μ l.

Step 4: Seal the tray. Make sure that the DNA-Mastermix-Taq solution has settled completely by gently tapping the tray on the working bench.

20 **Step 5:** Cycle on a thermocycler: 95°C 2min followed by 30 cycles of 94°C 20s, 63°C 20s, 72°C 1m30s.

Step 6: Separate the amplicons using 2% agarose gel electrophoresis of 8 μ l of the reaction at 150 V for 20-25 minutes.

 In all examples, the IHW KIR DNA samples were ordered
25 from IHWG (International Histocompatibility Working Group) at the Fred Hutchinson Cancer Research Center in Seattle, Washington. The primer sets used in the example are set forth in Table 2 or Table 3.

Example 1

As can be seen from Figure 2, twenty primer mixes identify the presence and absence of all presently known alleles of the KIR genes. The assay identifies 2DL1, 2DL2, 2DL3, 2DL4, 2DL5, 2DP1, 2DS1, 2DS2, 2DS3, 2DS4, 2DS5, 3DL1, 3DL2, 3DL3, 3DP1, and 3DS1 as measured against the sequence alignment resources found at IPD-KIR Sequence Database (<http://www.ebi.ac.uk/ipd/kir/>) and dbMHC (<http://www.ncbi.nlm.nih.gov/mhc/>). Each master mix contains a distinct internal control primer set to ensure proper assay performance (Tables 2 and 3 and Figures). A DNA size marker is used to demonstrate relative amplicon sizes. This assay discriminates the more recently described KIR variants 2DL5A and 2DL5B; 2DS4*00101/00101/002 and 2DS4*003; and 3DP1*001/002 and 3DP1*00301/00302. The allele information gained from the reaction may be used to deduce presently known KIR haplotypes.

Functionality of the primer mixes was challenged by testing with a range of variables similar to what may be encountered in routine laboratory use. Table 1 shows the parameters, ranges, and acceptance criteria used with the primer mixes. The parameters show a potential for conservation of sample and reagents.

Table 1

Validation Results		
Parameters	Criteria	
DNA template concentration	Higher end	No false positive
	Lower end	No false negative or failed reactions
Taq amount	Higher end	No false positive
	Lower end	No band dropout
Thermal cycler temperature	Denaturing temp. lower than 95°C	No band dropout
	Annealing temp. higher than 63°C	No band dropout
	Annealing temp. lower than 63°C	No false positive

The results discussed in Table 1 are shown in Figures 1 and 3-7. Lane assignments for these figures correspond to the wells and primers shown in Tables 2, 3 and/or Figure 2. Table 2 sets out the exact primer sequences used in the example. Positive results for a given allele are indicated by the presence of multiple bands per sample, as one band corresponds to an internal control. As can be seen from Figure 1, the results for two subsets of KIR DNA (IHW 1175 and IHW 1181) are consistent for all three thermal cyclers used. Figure 3 illustrates that different DNA template quantities, ranging from 15 ng to 250 ng all provided the same positive results. Figure 4 demonstrates that differing polymerase amounts also provided consistent allele identification. Figures 5-7 demonstrate successful allele identification at different thermocycling annealing and denaturing temperatures. These figures demonstrate that the present primer sets and methods are: specific as they produce the correct amplicon size, robust because specific and abundant amplicons exist at varied conditions, and sensitive relative to template amount.

Figure 8 shows selected IHW panel sample results. Figure 8 demonstrates that DNA samples encoding different KIR alleles can be successfully identified using the primers of Table 2.

In the above example, 800bp internal controls were used
5 (genbank acc#AF442818 C-reactive protein gene – the primer locations for the 800 bp internal control were CRP05 (5') - 18649-18667; CRP06 (3') - 19450-19430; CRP07 (5') - 18642-18663 and CRP08 (3') - 19448-19427), as well as 200bp internal controls (genbank acc# J04038 Human glyceraldehyde-3-phosphate dehydrogenase (GAPDH) gene – the primer
10 locations for the 200 bp internal control were 5'GPDH - 619-638; and 3'SGPDH - 815-796).

The present primers and kits can have any or all of the components described herein. Likewise, the present methods can be carried out by performing any of the steps described herein, either alone
15 or in various combinations. One skilled in the art will recognize that all embodiments are capable of use with all other appropriate embodiments described herein. Additionally, one skilled in the art will realize that certain embodiments also encompass variations of the present primers, configurations and methods that specifically exclude one or more of the
20 components or steps described herein.

As will be understood by one skilled in the art, for any and all purposes, particularly in terms of providing a written description, all ranges disclosed herein also encompass any and all possible subranges and combinations of subranges thereof. Any listed range can be easily
25 recognized as sufficiently describing and enabling the same range being broken down into at least equal halves, thirds, quarters, fifths, tenths, etc. As a non-limiting example, each range discussed herein can be readily broken down into a lower third, middle third and upper third, etc. As will also be understood by one skilled in the art all language such as “up to,”
30 “at least,” “greater than,” “less than,” “more than” and the like include

the number recited and refer to ranges which can be subsequently broken down into subranges as discussed above. In the same manner, all ratios disclosed herein also include all subratios falling within the broader ratio.

One skilled in the art will also readily recognize that where
5 members are grouped together in a common manner, such as in a Markush group, particular embodiments encompass not only the entire group listed as a whole, but each member of the group individually and all possible subgroups of the main group. Accordingly, for all purposes, certain embodiments encompass not only the main group, but also the
10 main group absent one or more of the group members. Individual embodiments also envisage the explicit exclusion of one or more of any of the group members.

All references, patents and publications disclosed herein are specifically incorporated by reference thereto. Unless otherwise specified,
15 "a" or "an" means "one or more."

While preferred embodiments have been illustrated and described, it should be understood that changes and modifications can be made therein in accordance with ordinary skill in the art without departing from the invention in its broader aspects as described herein. The broader
20 aspects of the present invention are defined in the following claims.

Table 2

Well #	KIR Allele Specificity	Sense Primer	Sense primer 3'end location	Antisense primer	Antisense primer 3'end location	Exon target	Internal Control size (bp)	App. PCR product size (bp)
1	2DL1*001-005	CATCAGTGGCATGACG	558	GGTCACTGGGAGCTGACAC	616	ex4	800	95
2	2DL2*001-004	AGAAACCTTCTCTCTCAGCCCA	686	GCCCTGCAGAGAACCTACA	790	ex5	800	145
3	2DL3*001-006	CTTCATCGCTGGTCTG	1094	CAGGCTCTTGGTCCATTACAA	1112	ex7-8	800	455
4	2DL4*00101/00102/00201/00202	GGTCTATATGAGAAACCTTCGCTTA	679	AGCCGAAGCATCTGTAGGTCT	866	ex5	800	230
5	2DL5A*001, 2DL5B*002-004	AGGTCTATTTGGGAAACCTTCA	675	ACTCATAGGGTGAGTCATGGAG	889	ex5	800	257
6	2DL5A*001	ACCATGTGCTCATGTGCA	15	CACAGGGCCCATGAGGAT	238	ex1-3	800	1753
7	2DL5B*002-003	CGTCACCTCCCATGATGTA	5'UT	CACAGGGCCCATGAGGAT	238	ex1-3	800	1893
8	2DS1*001-004	CTTCTCCATCAGTCGCATGAG	557	AGGGTCACTGGGAGCTGAC	616	ex4	800	140
9	2DS2*001-005	TGCACAGAGAGGGGAAGTA	482	CGGACACTCTCACCTGTGTG	648	ex4	800	207
10	2DS3*00101-00103	ACCTTGTCTTGCAGCTGCT	739	GAAGCATCTGTAGGTTCTCTCT	861	ex5	800	162
11	2DS4*00101/00102/002	CAGCTCCCGGAGCTCCTA	749	TGACGGAAACAAGCAGTGGA	927	ex5	800	215
12	2DS4*003 (KIR1D)	CCTTGTCTTGCAGCTCCATC	763	TGACGGAAACAAGCAGTGGA	927	ex5	800	200
13	2DS5*001-003	AGAGAGGGGACGTTTAACC	487	TCCAGAGGGTCACTGGGC	624	ex4	800	179
14	3DL1*00101/00102/002/03/00401/00402/005-009	TGAGCACTTCTTTCTGCACAA	470	GTAGGTCCTGCAGGKCAA	560	ex4	800	129
15	3DL2*001-012	AACCTTCTCTGTCTGCCC	100	GGAAGATGGGAACGTGGC	197	ex3	800	133
16	3DL3*001/00201/00202/03/004	CCTGCAATGTTGGTCAGATG	442	GAGCCGACAACCTCATAGGGTA	605	ex4	800	203
17	3DS1*010-014	CGCTGTGGTGCCTCGC	123	ACCTGTGACCATGATCACCAT	337	ex3	800	250
18	2DP1*001/002	ACATGTGATTCCTGGGTGCAT	150	TGTGAACCCCGACATCTGTAC	276	ex3	800	171
19	3DP1*001/002	CTTCCAGGGTCTCTTCTGCTGC	49	GAAAACGGGTTCGCGAATAC	223	ex2-3	200	975
20	3DP1*00301/00302	TCCGCTGCTGAGCTGAG	5'UT			ex1-3	200	344
	Negative Control						NONE	200
								800

Table 3

Well #	KIR Allele Specificity	Sense primer 3'end	Sense primer 3'end location	Antisense primer 3'end	Antisense primer 3'end location	App. Internal control size (bp)	App. PCR product size (bp)
1	2DL1*001-005	GAA	453	GCG	557	800	145
2	2DL2*001-004	CCA	686	ACA	790	800	145
3	2DL3*001-006	CTG	1094	CAA	1112	800	455
4	2DL4*00101/00102/00201/00202/003-007	TTA	679	TCT	866	800	230
5	2DL5A*001, 2DL5B*002-004	TCA	675	GAG	889	800	257
6	2DL6A*001	TCA	16	GAT	238	800	1753
7	2DL6B*002-004	GTA	5'UT	GAT	238	800	1893
8	2DS1*001	GAG	557	GAC	618	800	100
9	2DS2*001-005	GAA	557	ATG	648	800	207
10	2DS3*00101-00103	GTA	482	ATG	648	800	162
11	2DS4*00101/00102/002	CCT	739	GCT	861	800	215
12	2DS4*003	CTA	749	GGA	927	800	200
13	2DS5*001-003	ATC	763	GGA	927	800	179
14	3DL1*00101/00102/002/003/00401/00402/005-009	ACC	487	GGC	624	800	129
15	3DL2*001-012	CAA	470	CAA	560	800	133
16	3DL3*001/00201/00202/003/004	CCC	100	GGC	197	800	203
17	3DS1*010-014	ATG	442	GTA	605	800	260
18	2DP1*001/002	CGC	123	CAT	337	800	171
19	3DP1*001/002	CAT	150	TAC	276	800	975
20	3DP1*00307/00302	TGC	49	TAC	223	200	344
21	Negative Control	GAG	5'UT	TAC	223	200	975
		TGC	49	TAC	223	200	200
					NONE	NONE	800

The three nucleotide sequence given in the table is the sequence of the last three nucleotides of the sense and antisense primers which together determine the allele specificity of each primer mix. The 3' end sequence of the antisense primer should be read in reverse direction complementary to the sense sequence.

The location of the last base of each primer is given in the table. The numbers correspond to the nucleotide number, not the amino acid codon number. The location of the first nucleotide corresponds to the beginning of the first codon.